Information Extraction from Swedish Medical Prescriptions with Sig-Transformer Encoder

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Abstract

Relying on large pretrained language models such as Bidirectional Encoder Representations from Transformers (BERT) for encoding and adding a simple prediction layer has led to impressive performance in many clinical natural language processing (NLP) tasks. In this work, we present a novel extension to the Transformer architecture, by incorporating signature transform with the self-attention model. This architecture is added between embedding and prediction layers. Experiments on a new Swedish prescription data show the proposed architecture to be superior in two of the three information extraction tasks, comparing to baseline models. Finally, we evaluate two different embedding approaches between applying Multilingual BERT and translating the Swedish text to English then encode with a BERT model pretrained on clinical notes.

1 Introduction

Medical prescription notes written by clinicians about patients contains valuable information that the structured part of electronic health records (EHRs) does not have. Manually extracting and annotating such information from large amount of textual fields in medical records such as prescription notes is time-consuming, and expensive since it has to be provided by domain specialists who are in high demand. Information extraction (IE), a specialised area in natural language processing (NLP), refers to the automatic extraction of structured information such as concepts, entities and events from free text (Wang et al., 2018). While rulebase IE systems are still widely used in healthcare (Banda et al., 2018; Davenport and Kalakota, 2019), machine learning-based data-driven approaches have gained much more interest due to their ability to scale and learn to recognise complex patterns. In this paper, we present a set of medical prescriptions

written in Swedish and develop a neural networkbased system that extracts important information from the textual data.

Recent advances in pretraining large-scale contextualised language representations, including ELMo (Peters et al., 2018) and BERT (Devlin et al., 2018), have proven to be a successful strategy for transfer learning and pushed the performance in many NLP tasks of general purpose. Clinical notes differ substantially to general-domain text and biomedical literature, in terms of its linguistic characteristics. Several studies have finetuned general-domain language models such as BERT on in-domain clinical text (e.g. electronic health records) (Si et al., 2019; Peng et al., 2019; Alsentzer et al., 2019; Huang et al., 2020) for downstream clinical NLP tasks. As summarised in (Gu et al., 2020) many such tasks can be formulated as a classification or regression task, wherein either a simple linear layer or sequential models such as LSTM or conditional random field (CRF) are added after the BERT encoding of the input text as task-specific prediction layer.

BERT's model architecture is based on the Transformer model (Vaswani et al., 2017), which employs the self-attention mechanism to attend to different positions of the input sequence. As it doesn't contain any recurrence or convolution, the model has to add positional encoding in order to model the order of the sequence (e.g. word order). Signature transform, initially introduced in rough path theory as a branch of stochastic analysis, is a non-parametric approach of encoding sequential data while capturing the order information in the data. It has shown successes in the recent years in a range of machine learning tasks involving sequential modelling (Arribas et al., 2018; Morrill et al., 2020; Toth and Oberhauser, 2020; Wang et al., 2020). We hypothesise the signature transform method can be integrated in the Transformer

model, in which its ability to naturally capture sequential ordering complements the ability of attending to important positions and learning long-range dependencies in self-attention.

As a preliminary study, our aim is to develop a machine learning-based system that extracts relevant information from the Swedish medical prescription notes¹, namely *quantity*, *quantity tag* and *indication*. The contributions of this work are as follows:

- We experiment two different approaches of embedding the Swedish prescription notes. One encodes the Swedish text directly using Multilingual BERT (M-BERT) (Devlin et al., 2018) while for the other approach we translate the prescriptions and then apply Clinical-BERT (Huang et al., 2020) that is pretrained on clinical text in English.
- 2. We propose an extension to the Transformer model, named Sig-Transformer Encoder (STE), which integrates signature transform into the Transformer architecture so the order information in the prescription notes can be learnt in a more effective way. To the best of our knowledge, this is the first attempt to integrate signature transform in the Transformer architecture.
- 3. We demonstrate good performance in two of the three tasks, namely *quantity* and *quantity tag*. As for the *indication* task, we provide an analysis on why the models fail.
- 4. We show one of our proposed STE-based approaches, namely M-BERT+STE, outperforming other baseline models.

2 Background

Clinical language representation has attracted an increasing interest in the natural language processing (NLP) community (Kalyan and Sangeetha, 2020). More recently with the advent of Bidirectional Encoder Representations from Transformers (BERT) (Devlin et al., 2018), fine-tuning of general-domain language models has been widely adopted for many clinical NLP tasks². Among them two clinicalBERT studies (Alsentzer et al., 2019; Huang et al., 2020) have conducted fine-tuning from either BERT or BioBERT (Lee et al., 2020) using de-identified clinical notes from MIMIC-III (Johnson et al., 2016). In particular, Huang et al. (2020) showed the effectiveness of its ClinicalBERT-based model for predicting hospital readmissions.

Many NLP applications in the medical domain can be formulated as either token classification, sequence classification or sequence regression, in which a pretrained language model such as clinicalBERT can be used to encode the input token sequence (which returns either the encoding for every input token or just the [CLS] token) and then a task-specific prediction model is added on top to generate the final output (Gu et al., 2020). In this section, we describe the Transformer encoder and the signature transform, which are the bases of our proposed architecture.

2.1 Transformer Encoder

The Transformer model employs an encoderdecoder structure (Vaswani et al., 2017). Its encoder, which is the model architecture of BERT, is composed of a stack of N identical layers. Each layer has a multi-head self-attention mechanism followed by a position-wise fully connected feedforward network. The attention function used in the Transformer model takes three input vectors: query (Q), key (K) and value (V). It generates an output vector by computing the weighted sum of the values. The weights are computed by the dot products of the query and all keys, scaled and applied a softmax function.

$$\operatorname{Attention}(Q, K, V) = \operatorname{softmax}(\frac{QK^T}{\sqrt{d_k}})V \quad (1)$$

Multi-head attention splits Q, K and V into multiple heads by linearly projection, which allows the model to jointly attend to information at different positions from different representation subspaces. Each projected head goes through the scaled dotproduct attention function, then concatenated and projected again to output the final values. The Transformer encoder does not explicitly model position information in its structure, and instead it requires adding representations of absolute positions (i.e. positional encoding) to its inputs. Shaw et al. (2018) presented an extension to self-attention, which incorporates relative position information

¹Our ultimate goal is to apply such model to a much larger database. However, this is beyond the scope of this paper.

²Gu et al. (2020) have challenged this strategy and reported better performance for PubMed-based biomedical tasks by conducting domain-specific pretraining from scratch. How-

ever, it did not address the clinical domain and has left it for future work.

for the input data. In this work, we propose to integrate signature transform with self-attention given that path signatures have been proven to be an effective way of capturing the sequential order information in the data.

2.2 Signature Transform

The theory of rough paths, developed by Lyons (1998), can be thought of as a non-linear extension of the classical theory of controlled differential equations. *Path signature* or simply *signature*, is an infinite collection of statistics characterising the underlying path (a discretised version of a continuous path), and *signature transform* is the map from a path to its signature³. Consider a *d*-dimensional time-dependent path *P* over the time interval $[0,T] \subset \mathbb{R}$, to a continuous map $P : [0,T] \to \mathbb{R}^d$. The signature S(P) of this path *P* over time interval [0,T] is the infinite collection of all iterated integrals of *P* such that every continuous function of the path may be approximated arbitrarily well by a linear function of its signature:

$$S(P)_{0,T} = (1, S(P)^{1}_{0,T}, \dots, S(P)^{d}_{0,T}, S(P)^{1,1}_{0,T}, \dots)$$

where the 0^{th} term is 1 by convention, and the superscripts of the terms after the 0^{th} term run along the set of all multi-index $\{(i_1, \ldots, i_k) | k \ge 1, i_1, \ldots, i_k \in [d]\}$ with the coordinate iterated integral being:

$$S(P)_{0,T}^{i_1,\ldots,i_k} = \int_{\substack{t_1 < \cdots < t_k \\ t_1,\ldots,t_k \in [0,T]}} dP_{t_1}^{i_1} \otimes \cdots \otimes dP_{t_k}^{i_k}$$

where $\forall k \geq 1$, $P_t \in \mathbb{R}^d$, $\forall t \in [0, T]$. $S(P)_{0,T}^{i_1, \dots, i_k}$ is termed as the *k*th level of the signature. In practice we truncate the signature to order *n*, where the degree of its iterated integrals is no greater than *n*. This ensures the path signature has finite dimensional representation. Let $TS(P)_{0,T}^n$ denote the truncated signature of *P* of order *n*, i.e.

$$TS(P)_{0,T}^n = (1, S(P)_{0,T}^1, \dots, S(P)_{0,T}^{k_n})$$

Therefore the dimensionality of the truncated path signature is $(d^{n+1} - d)(d - 1)^{-1}$. We describe signature transform in more (mathematical) details in Appendix A.2.

In recent years using path signatures as features in a suitable neural network model has shown success in various applications, such as online handwritten Chinese character recognition (Yang et al., 2016; Xie et al., 2018), action recognition in videos (Yang et al., 2017) and speech emotion recognition (Wang et al., 2019). More recently, Kidger et al. (2019) proposed to use signature transform deeper within a network, rather than as a feature transformation. Kidger and Lyons (2020) developed differentiable computation of signature transform on both CPU and GPU. To the best of our knowledge, we are the first to incorporate signature transform in the Transformer architecture.



Figure 1: The Sig-Transformer Encoder. d_{sig} is the dimension size in the truncated signature, and d_{model} is the dimension size in the embedding layer as well as before and after the second feed-forward sub-layer.

3 Sig-Transformer Encoder

In this section, we present our proposed extension to the Transformer Encoder (TE), named Sig-Transformer Encoder (STE). As depicted in Figure 1, STE consists of a stack of N identical layers, and each layer has three sub-layers. It replaces the multi-head self-attention sub-layer in TE with our Additive Multi-Head Sig-Attention mechanism, followed by two position-wise fully connected feed-forward sub-layers. Same as the Transformer model, we also employ a residual connection (He et al., 2016) at the last sub-layer. followed by layer normalisation (Ba et al., 2016). The two feed-forward sub-layers and the embed-

³We refer the reader to (Lyons, 2014) for a rigorous introduction of signature transform, and (Chevyrev and Kormilitzin, 2016) for a primer on its use in machine learning.



Figure 2: (left) Sig-Attention mechanism; (right) Additive Multi-Head Sig-Attention.

ding layers, produce output of the same dimension $d_{model} = 768$.

3.1 Sig-Attention

The core idea behind our proposed Additive Multi-Head Sig-Attention is the Sig-Attention mechanism (Figure 2 left). It takes the query (Q), key (K) and value (V) vectors as input, and performs three consecutive transformations: (1), scaled dot-product attention; (2), position-wise fully linear map, for dimensionality reduction; (3), signature transform. Its output is computed as follows:

> SigAttention(Q, K, V) =ReducedSig(Attention(Q, K, V))

where

- Attention(Q, K, V) is as defined in Equation (1);
- Reduced Sig $(x) = S^N(xW + b)$ for any x;
- S^N is the signature truncated at a given order N;
- W and b are respectively the weight and bias matrices of the linear map.

As the size of the truncated signature increases exponentially with its input dimension, it is important to reduce the size of attended context vector before applying signature transform. We refer the readers to Table 5 for examples of how the size of the truncated signature (d_{sig}) is determined by its input dimension (d_{presig}) and the order of signature truncation $(order_{sig})$.

3.2 Additive Multi-Head Sig-Attention

In the proposed Additive Multi-Head Sig-Attention model, we combine the information from the input sequence with the output of Sig-Attention, in different representation subspaces (Figure 2 right). The model takes the embedding of an input sequence X as well as the queries, keys and values as input:

$$\begin{split} \text{AdditiveMultiHead}(X,Q,K,V) = \\ & \text{Concat}(\text{head}_1,...,\text{head}_h) \\ \text{where head}_i = \text{ReducedSig}(X) + \\ & \text{SigAttention}(QW_i^Q,KW_i^K,VW_i^V) \end{split}$$

and h is the number of parallel heads, the projections are parameter matrices $W_i^Q \in \mathbb{R}^{d_{\text{model}} \times d_k}$, $W_i^K \in \mathbb{R}^{d_{\text{model}} \times d_k}$, $W_i^V \in \mathbb{R}^{d_{\text{model}} \times d_k}$ and $d_k = d_{model}/h$.

As shown in Figure 2 (right), we linearly project the queries, keys and values as well as the input embeddings to d_{model} and d_{presig} dimensions. Then the query, key and value vectors will be fed into the Sig-Attention mechanism, while having been reduced its dimensions by the linear projection the input vector X will be applied to signature transform. The two output signatures are added, resulting in one signature vector $s_i \in \mathbb{R}^{d_{sig}}$ for each head *i*. Finally all the signature vectors are concatenated and passed onto the next sub-layer.

3.3 Position-wise Feed-Forward Networks

The second sub-layer in Figure 1 is a linear transformation $W \in \mathbb{R}^{d_{sig} \times d_{model}}$ which is applied to each position separately and identically. It simply brings back the dimensions from d_{sig} to d_{model} . The third sub-layer is a ReLU activation followed by another linear transformation. The dimensionality of its input and output is d_{model} .

3.4 Dropout

Dropout is applied to the output of the three sublayers as well as after the ReLU activation in the third sub-layer. We use a rate of $P_{drop} = 0.1$, as is used in (Vaswani et al., 2017). Moreover, residual connection and layer normalisation are used only in the third sub-layer.

4 Data

Our proposed Sig-Transformer Encoder can be used for general purpose representation learning, and any classifier or regressor can be added on top of it to perform down-stream tasks. We conduct all of our experiments with a medical prescription data for the class of "beta-blockers (ATC code C07)", provided by the Karolinska University Hospital. It is sampled from a much larger database of 41 million medical prescription records. This dataset contains 3,852 non-duplicated prescriptions written in Swedish. Some examples of annotated prescriptions are shown in Table 1.

The medical practitioners at the hospital have annotated three labels, in which we use QUANTITY for regression while QUANTITY TAG and INDICA-TION as two classification tasks:

- QUANTITY: the total amount of tablets or capsules prescribed. The values are multiples of 0.5;
- QUANTITY TAG (5 classes): the label or tag of the quantity prescribed to the patient;
 - 1. *Not Specified*: the quantity was not specified in the prescription;
 - 2. *Complex* : a range of quantities was given. In that case, the quantity is an average between the minimum and the maximum quantities;
 - 3. *PRN* : prescription to take only if needed;
 - 4. *As Per Previous Prescription*: refers to guidance in previous prescription;
 - 5. Standard: standard prescription;
- INDICATION (5 classes): the purpose of the prescription. It originally had 44 classes where many account for only one record⁴. The

medical practitioners have aggregated them into 5 medically meaningful classes: *Cardiac*, *Tremors*, *Migraine*, *Others*, and *NA* (Not Annotated).

One challenge of the data is the remarkable class imbalance in its labels, as shown in Table 2. It's still prominent in the INDICATION field after aggregating the original classes, where *Cardiac* and *NA* account for majority of the instances, while the other 3 classes are seldom annotated in the data. Another challenge, which is prevalent in electronic health record (EHR) datasets, is the free-form nature of its text and the writing styles vary considerably between different doctors.



Figure 3: Multi-task learning architecture without Sig-Transformer Encoder (STE).



Figure 4: Our proposed multi-task learning architecture with Sig-Transformer Encoder (STE).

4.1 Data Preprocessing

As mentioned in Section 1 we experiment two different embedding models for the Swedish prescription records, in which one of them, namely Clinical-BERT (Huang et al., 2020), requires its input text to be English. We use the Google Translate API to obtain translated English prescriptions. Minimal text preprocessing was taken, and all prescription text including non-ASCII characters (e.g. å, ä, ö) was read with UTF-8 encoding.

⁴The original 44 classes and the number of prescriptions per class, can be found in Figure 5.

Swedish	Translated English	Indication	Quantity	Quantity Tag
1 TABLETT FOREBYGGANDE	1 tablet prevention	Migraine	ina 1	Standard
MOT MIGRAN	against migrain	wingrame	1	Stanuaru
MOT HOGT BLODTRYCK	Against high blood pressure			
OCHHJARTKLAPPNING	and heart palpitations	Cardiac-	2	Standard
2 TABLETTER KLOCKGAN 08:00	2 tablets at 08:00,	hypertension	5	Standard
1 TABLETT KLOCKAN 18:00.	1 tablet at 18:00.			
2 TABLETTER KL. 08,	2 tablets kl. 08,			Standard
2 TABLETT KL. 20. DAGLIGEN.	2 tablet kl. 20. Daily.	NA	4	
OBS KVALLSDOSEN HAR	Note the evening box has		4	Standard
HOJTS JFRT MED 070427.	hojts jfrt with 070427.			
1 tablett vid behov mot stress	1 tablet if needed against stress	Anxiety	1	PRN
FOR BLODTRVCK	For blood pressure	Cardiac-		
OCH HIADTDYTM	and heart rhythm	hypertension-	0	Not Specified
OCHTIJAKTKI IWI-	and neart my thin -	dysrhythmia		
1 tabl pA morgonen och	1 table in the morning and	Cardiaa		
en halv tabl pA kvAllen	a half table in the evening	Cardiac-	1.5	Standard
fOr hjArtrytmen.	for the heart rhythem.	uysinyuiiiia		
1-2 tablett 2 gAnger dagligen	1-2 tablets 2 times daily	NA	3	Complex

Table 1: Example prescriptions with translations and annotations. The second column is the English translation obtained from Google Translate API. The last three columns are the labels of interest which we aim to extract automatically. Some longer example prescriptions can be viewed in Table 8.

QUANTITY TAG			INDICATION			
Standard	3489	90.6%	Cardiac	2980	77.4%	
PRN	89	2.3%	Tremors	81	2.1%	
APPP	40	1.0%	Migraine	69	1.8%	
Complex	29	0.8%	Other	15	0.4%	
NS	205	5.3%	NA	707	18.4%	

Table 2: Class distribution for QUANTITY TAG and IN-DICATION. APPP is an abbreviation for *As Per Previous Prescription*, NS stands for *Not Specified* and NA stands for *Not Annotated*.

5 Multi-Task Learning

Our objective is to automatically find the QUAN-TITY of prescribed medicine as well as QUANTITY TAG and INDICATION of the prescription, for each prescription note. We formulate this as a multi-task learning problem consisting of a regression task and two classification tasks. As depicted in Figure 3, such multi-task learning model uses BERT or any of its variants to encode each input prescription note represented by the respective [CLS] token, then we can add three separate predictors on top of it and all three jointly learn with the embedding layer. We use this architecture as one of our baseline approaches.

In our proposed multi-task learning architecture, we add Sig-Transformer Encoder (STE) in-between the embedding layer (BERT) and the three predictors in order to learn the sequential order information in the input text encoded by the token-level BERT representations (instead from [CLS]). The three predictors and STE are jointly trained. We compare the two architectures and other baseline models in Section 6.

6 Experiments

6.1 Experiment Setup

As described in Section 2, BERT and its variants have been used in a range of clinical machine learning tasks. Considering our data is in Swedish we explore two different approaches for encoding the prescription notes: (1), Apply Multilingual BERT (M-BERT) (Devlin et al., 2018) directly to the Swedish text; (2), Translate the prescriptions to English as described in Section 4.1, and then encode the translated text with ClinicalBERT (Huang et al., 2020). ClinicalBERT is fine-tuned on clinical notes, therefore it has embedded more domain knowledge and has proven to be more effective over BERT-base on several clinical tasks. However, any translation error would affect the performance of subsequent models.

We use the PyTorch implementation of Transformers by Hugging Face⁵ for loading M-BERT and ClinicalBERT, and *Signatory*⁶ (Kidger and Lyons, 2020) for differentiable computations of

⁵https://github.com/huggingface/transformers

⁶https://github.com/patrick-kidger/signatory

signatures on GPU. We use a linear layer as the classifier for QUANTITY TAG, and two-layer network for QUANTITY and INDICATION respectively. More details of the implementation choices are described in Appendix A.1. We use cross-entropy (CE) loss for the two classification tasks and meansquare error (MSE) for the regression task. The overall loss function is a weighted average of the loss functions of all three tasks:

 $\mathcal{L} =$

 $\alpha_{qnt} \times MSE_{qnt} + \beta_{qntt} \times CE_{qntt} + \beta_{ind} \times CE_{ind}$

where *qnt* stands for quantity, *qntt* stands for quantity tag, *ind* stands for indication, and $\alpha_{qnt} + \beta_{qntt} + \beta_{ind} = 1$. α_{qnt} , β_{qntt} and β_{ind} are parameters to be optimised.

To compare model performance we use the simple multi-task learning models described in Section 5 without adding STE (Figure 3), as the baseline systems. We also replace STE with LSTM for comparison as both learn sequential information from the input data⁷. All experiments are conducted using 5-fold cross validation, where we use 3 folds for training, 1 fold for validation and 1 fold as the test set at each iteration. We repeat the experiments for each model with 10 different random seeds and average the scores as the final result.

6.2 Results

The results for all three tasks are summarised in Table 3. We first notice all models have failed to recognise the INDICATION classes, which we will discuss in Section 6.3. In this section we mainly discuss the performance for QUANTITY and QUANTITY TAG.

Among the three models without the use of LSTM or STE, both ClinicalBERT and Multilingual BERT (M-BERT) outperform the BERT-base model in all three tasks. As for the three LSTMbased models, using the BERT-base representations achieves the best performance while ClinicalBERT and M-BERT are comparable with each other. Among our proposed STE-based models, M-BERT+STE outperforms the other two variants.

Comparing the results across the board, adding LSTM or STE improves the model performance in most cases except for ClinicalBERT, where the

addition of LSTM or STE has worsened the performance for ClinicalBERT. We think the potential discrepancies in the translation of the Swedish prescriptions have possibly affected the working of LSTM and STE, as both models take all the translated token embeddings of each prescription as input where the simple ClinicalBERT baseline only takes the encoding of the [CLS] token as input. Comparing between the two different prescription embedding approaches, translating them then apply ClinicalBERT clearly beats using the multilingual-BERT when LSTM or STE is not added. When we incorporate LSTM or STE in the model, the multilingual embedding approach has obtained better results in most cases. Overall, one of our proposed models, M-BERT+STE, achieves the best performance for the QUANTITY and QUANTITY TAG tasks⁸.

In Table 6 we present results obtained by each model across different classes of QUANTITY TAG. We can see the *Standard* class (i.e. standard prescription) followed by *PRN* (i.e. prescription to take only if needed) and *APPP* (i.e. as per previous prescription) have received better classifications by the three STE models. The models also performed well in the *Not Specified* class. The *Complex* class refers to a range of quantities given in the prescription. As expected, many models do not perform well in this class. The overall best performing model, M-BERT+STE, also performs consistently well across different classes of QUANTITY TAG.

6.3 The difficult case of Indication

The class imbalance is remarkable for QUANTITY TAG and INDICATION in this dataset, as described in Section 4. The strong results obtained from many models for QUANTITY TAG suggests albeit the class imbalance the boundaries between its classes are distinguishable and can be learnt from fewer examples. INDICATION is a more challenging task. The aggregation of the original 44 classes to the final 5 classes has reduced the severity of the class imbalance issue⁹. However, the new classification becomes much less distinguishable and the new classes are less distinct as each class now contains

⁷Unlike the Transformer model, recurrent neural networks (RNNs) such as LSTM, model relative and absolute positions along the time dimension directly through their sequential structure (Shaw et al., 2018).

⁸The results obtained by M-BERT+STE and ClinicalBERT are different at the significance level of 0.1, with p = 0.068and 0.065 for QUANTITY and QUANTITY TAG respectively, using Wilcoxon signed-rank test.

⁹We have experimented with the original 44 classes, and obtained terrible results. We have also tried setting class weights to alleviate the class imbalance issue with no significant difference observed in the final results.

Model	QUANTITY	QUANTITY TAG	INDICATION
Base	0.50	0.60	0.08
ClinicalBERT	0.21	0.89	0.09
M-BERT	0.41	0.63	0.10
Base + LSTM	0.45	0.76	0.06
ClinicalBERT + LSTM	0.47	0.64	0.10
M-BERT + LSTM	0.50	0.68	0.02
Base + STE	0.23	0.78	0.03
ClinicalBERT + STE	0.36	0.77	0.06
M-BERT + STE	0.15	0.92	0.05

Table 3: Performance comparison between our proposed STE-based approaches and baseline models. The QUAN-TITY task is measured in mean squared error (MSE), while for QUANTITY TAG and INDICATION we use Macro F-1 score. Base refers the original BERT-base model, M-BERT is the Multilingual BERT model pretrained on Swedish text data. STE refers to Sig-Transformer Encoder.

a range of topics, as shown in Figure 5. Such low scores are also observed across different classes of INDICATION in Table 7.

6.4 Ablation Study

The positional encoding proposed in (Vaswani et al., 2017) is based on sinusoids of varying frequency, and the authors hypothesise this would help the model to generalise to sequences of variable lengths during training. Our proposed architecture has both the absolute positional representations of words and the signature transform that is also invariant to the sequence length for encoding order of events. Here we conduct ablation study by removing the positional encoding or the signature transform components from the best performing model, M-BERT+STE, and compare the results. As shown in Table 4, we obtain worse performance without positional encoding or without signature transform, for all three tasks, suggesting the contributions of using the absolute positional encoding and the proposed integration of signature transform.

Model	Quantity	Quantity Tag	Indication
Full model	0.15	0.92	0.05
w/o PE	0.38	0.81	0.01
w/o ST	0.31	0.77	0.02

Table 4: Ablation study of the best performing M-BERT + STE model, removing positional encoding (PE) or signature transform (ST).

7 Conclusions and Future Work

In this work, we propose a new extension to the Transformer architecture, named Sig-Transformer Encoder (STE), by incorporating signature transform with the self-attention mechanism. As a preliminary study, we aimed to automatically extract information related to *quantity*, *quantity tag* and the *indication* label from a Swedish medical prescription dataset. We demonstrated good performance in two of the three tasks in a multi-task learning framework. Lastly, we compared two embedding approaches between applying Clinical-BERT (on translated text) and Multilingual BERT (M-BERT). Although one of our proposed models, namely M-BERT+STE, reported the best performance for *quantity* and *quantity tag*, all models failed to perform on the *indication* task which we provided possible explanations.

For future work, we plan to apply the proposed STE models to the much larger prescription database and investigate ways to improve the labelling for *indication*. We also plan to benchmark our models on other clinical tasks such as predicting hospital readmission (Huang et al., 2020), as well as the more general NLP tasks evaluated in (Devlin et al., 2018). Moreover, a further investigation to better understand the contribution of signature transform and the interplay between signature and attentions would be very insightful.

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A Appendices

A.1 Implementation Details

Our classifier for INDICATION is a two-layer network, in which the first layer has the size of 50 and 5 for the second layer. ReLU activation and dropout are used in-between the two layers. The dropout rate is set to be 0.1. We also use a twolayer network for the QUANTITY regression task, in which the first layer has the size of 10 and its second layer predicts the value for *quantity*. We also add a dropout the rate 0.1 in-between the two layers. Our classifier for QUANTITY TAG has only one layer the size of 5.

We reduce the dimension of attended vector before signature transform from 768 to 32, as depicted in Figure 2 (left). We set the order of truncated signature to be 2, and employ 8 parallel attention layers (or heads). We ran grid-search over two choices of learning rate lr : [0.00003, 0.00005], and 10 randomly sampled triplets of α_{qnt} , β_{qntt} and β_{ind} . Each value of α_{qnt} , β_{qntt} and β_{ind} is between 0 and 1 respectively, while we make sure each combination of the three values sums to 1.

A.2 Signature of paths

We begin with the definition of the signature, using more traditional notation of stochastic calculus.

Definition 1. Let $X = (X^1, ..., X^d)$ be a path in \mathbb{R}^d . The signature of X is defined as the infinite collection of iterated integrals:

$$S(X) = \left(\int \dots \int_{a < t_1 < \dots < t_k < b} dX_{t_1} \otimes \dots \otimes dX_{t_k}\right)_{k \ge 0}$$
$$= \left(\left(\int \dots \int_{a < t_1 < \dots < t_k < b} dX_{t_1}^{i_1} \dots dX_{t_k}^{i_k}\right)_{1 \le i_1, \dots, i_k \le d}\right)_{k \ge 0}$$

where $dX_t = \frac{dX_t}{dt} dt$ and the k = 0 term is taken to be $l \in \mathbb{R}$.

Definition 2. *The truncated signature of order N of X is defined as:*

$$S^N(X) = \left(\int \dots \int_{a < t_1 < \dots < t_k < b} dX_{t_1} \otimes \dots \otimes dX_{t_k}\right)_{0 \le k \le N}.$$

The dimension of the truncated signature explodes exponentially with the input path dimension:

Proposition 1. For any $d \ge 1$, the truncated signature of order N of a d-dimensional path has the dimension of:

$$\sum_{k=0}^{N} d^k = \frac{d^{N+1} - 1}{d-1}$$

In practice, the term of order 0 is dropped as it is always equal to 1. For clarity, we define:

$$S(X)^{i_1,...,i_k} = \int \dots \int_{a < t_1 < \ldots < t_k < b} dX^{i_1}_{t_1} \dots dX^{i_k}_{t_k}$$

with $1 \leq i_1, ..., i_k \leq d$, so that:

$$S(X) = \left(\left(S(X)^{i_1, \dots, i_k} \right)_{1 \le i_1, \dots, i_k \le d} \right) \right)_{k \ge 0}$$

= $(1, S(X)^1, \dots, S(X)^d, S(X)^{1,1}, S(X)^{1,2}, \dots).$

d_{presig}	$\operatorname{order}_{sig}$	d_{sig}	d_{presig}	$\operatorname{order}_{sig}$	d_{sig}
512	1	512	16	3	4K
512	2	262K	8	2	72
256	2	66K	8	3	584
128	2	16K	8	4	5K
128	3	2M	4	4	340
64	2	4K	4	5	1365
64	3	266K	4	6	5K
32	2	1057	2	9	1022
32	3	34K	2	10	2K
16	2	272	2	12	8K

Table 5: The number of dimensions (d_{sig}) of the truncated signature is determined by the size of its input (d_{presig}) and the order of truncation selected (order_{sig}).

Madal						
WIOUEI	Standard	APPP	PRN	Complex	NS	QUANTITY
Base	0.71	0.50	0.10	0.76	0.95	0.50
ClinicalBERT	0.83	0.97	0.89	0.99	0.79	0.21
M-BERT	0.94	0.41	0.04	0.81	0.98	0.41
Base + LSTM	0.93	0.36	0.99	0.77	0.74	0.45
ClinicalBERT + LSTM	0.90	0.23	0.99	0.61	0.49	0.47
M-BERT + LSTM	0.29	0.99	0.72	0.47	0.93	0.50
Base + STE	0.97	0.84	0.77	0.44	0.88	0.23
ClinicalBERT + STE	0.99	0.70	0.85	0.65	0.65	0.36
M-BERT + STE	0.86	0.97	1.00	0.89	0.86	0.15

Table 6: Model performance comparison for QUANTITY and also across different classes in QUANTITY TAG. APPP: As Per Previous Prescription; NS: Not Specified.

Madal	INDICATION							
wiouei	Cardiac	Tremors	Migraine	Others	NA			
Base	0.03	0.05	0.02	0.27	0.03			
ClinicalBERT	0.00	0.24	0.03	0.04	0.13			
M-BERT	0.05	0.01	0.09	0.35	0.00			
Base + LSTM	0.02	0.00	0.00	0.31	0.00			
ClinicalBERT + LSTM	0.01	0.03	0.34	0.08	0.02			
M-BERT + LSTM	0.05	0.00	0.00	0.04	0.03			
Base + STE	0.00	0.00	0.02	0.08	0.05			
ClinicalBERT + STE	0.00	0.26	0.05	0.02	0.00			
M-BERT + STE	0.00	0.11	0.00	0.00	0.15			

Table 7: Model performance comparison across different classes in INDICATION. NA: Not Annotated.

Indic	ation (4	44)	
CARDIAC - HYPERTENSION - PALPITATIONS	49		
CARDIAC - PALPITATIONS - TREMORS	1	ESOPHAGEAL VARICES	1
CARDIAC - PALPITATIONS - ANGINA	1	ANXIETY - CARDIAC - PALPITATIONS	3
CARDIAC - PALPITATIONS - ATRIAL FIBRILLATION	1	ANXIETY	6
CARDIAC - ATRIAL FIBRILLATION	57	ANXIETY - MIGRAINE	1
CARDIAC - DYSRHYTHMIA	101	HYPERTHYROIDISM	1
CARDIAC - HEART FAILURE	60	HYPERGLYCEMIA	2
CARDIAC - HYPERTENSION - ANGINA - PALPITATIONS	1	CHEST PAIN	1
CARDIAC - ATRIAL FIBRILLATION - DYSRHYTHMIA/RATE	1		
CARDIAC - HYPERTENSION - DYSRHYTHMIA	19	TREMORS	77
CARDIAC - MIGRAINE	1	TREMORS - CARDIAC - HYPERTENSION	1
CARDIAC - DYSRHYTHMIA/RATE	114	TREMORS - ANXIETY	1
CARDIAC - HYPERTENSION - ESOPHAGEAL VARICES	1	TREMORS - CARDIAC - PALPITATIONS	2
CARDIAC - PALPITATIONS	162		
CARDIAC - HYPERTENSION - HEART FAILURE	4		
CARDIAC - HYPERTENSION - MIGRAINE	12	MIGRAINE	69
CARDIAC - HYPERTENSION - ANGINA	28		
CARDIAC - TREMORS	2		
CARDIAC - DYSRHYTHMIA - MIGRAINE	1	NA	707
CARDIAC - ANGINA - ATRIAL FIBRILLATION	1		
CARDIAC - HEPATIC CIRCULATION	1		
CARDIAC - ANGINA	34		
CARDIAC - DYSRHYTHMIA/RATE - PALPITATIONS	1		
CARDIAC - HYPERTENSION	1655		
CARDIAC - HYPERTENSION - ATRIAL FIBRILLATION	2		
CARDIAC - ESOPHAGEAL VARICES	2		
CARDIAC	608		
CARDIAC - ANGINA - DYSRHYTHMIA/RATE	1		
CARDIAC - HYPERTENSION - DYSRHYTHMIA/RATE	50		
CARDIAC - PALPITATIONS - MIGRAINE	1		
CARDIAC - HYPERTENSION - TREMORS	8		

Figure 5: Number of prescriptions per indication class, where the indication label has the original 44 classes. *NA* stands for *Not Annotated*.

Quantity Tag	Standard	Complex	Standard	Standard	Standard	Standard
Quantity			1	1.5	1	c
Indication	Cardiac- hypertension	Cardiac- palpitations	NA	NA	Cardiac- hypertension	Cardiac- hypertension
Translated English	0.5 tablets 2 times daily. For the heart rhythm and reduces blood pressure The tablets are swallowed whole (can be divided at swallowing durations, but don't k	I table 1 time daily for 3 weeks. Successful increase for 2 tablets daily dependent on control and symptoms relief, against palpitations	1 tbale 1 time daily. Previously opening to a tablet morning and lunch but begin with a tablet early morning, against shakes	1/2 table on morning, 1/2 table on lunch and 1/2 table on evening against tremor, blood pressure control	1 tablet 1 time daily against high blood pressure you should order the painting lighting may june for control BT diabetes	2 prolonged-release tablets at. 08,1 prolonged-release tablet at. 20. Daily.Against high blood pressure
Swedish	0.5 TABLETTER 2 GANGER DAGLIGEN. FOR HJARTRYTMEN OCH SANKER BLODTRYCKET TABLETTERNA SVALJES HELA (KAN DELAS VID SVALJSVARIGHETER, MEN FAR EJ K	1 TABLETT 1 GANG DAGLIGEN I 3 VECKOR. EVENTUELL HOJNING TILL 2 TABLETTER DAGLIGEN BEROENDE PA KONTROLLEN OCH SYMTOMLINDRING, MOT HJARTKLAPPNING	1 TABLETT 1 GANG DAGLIGEN. EFTERHAND EVENTUELL OKNING TILL EN TABLETT MORGON OCH LUNCH MEN BORJA MED EN TABLETT TIDIG MORGON, MOT SKAKNINGAR.	1/2 TABLETT PA MORGONEN , 1/2 TABLETT PA LUNCHEN OCH 1/2 TABLETT PA KVALLEN MOT TREMOR, BLODTRYCKSREGLERANDE	1 TABLETT 1 GANG DAGLIGEN MOT HOGT BLODTRYCK DU BOR BESTALLA LAKARTIDLAMPLIGEN MAJ JUNI FOR KONTROLL BT DIABETES.	2 depottabletter kl. 08, 1 depottablett kl. 20. Dagligen. Mot hOgt blodtryck

Table 8: Longer example prescriptions with translations and annotations.